Amendments to the Specification:

Please replace the paragraph beginning on page 54, line 22, with the following paragraph.

In one embodiment, suitable compounds include, but are not limited to, verapamil, desmethoxyverapamil, chloroquine, quinine, chinchonidine, primaquine, tamoxifen, dihydrocyclosporin, yohimbine, corynanthine, reserpine, physostigmine, acridine, acridine orange, quinacrine, trifluoroperazine chlorpromazine, propanolol, atropine, tryptamine, forskolin, 1,9-dideoxyforskolin, cyclosporin, (US Patent 4,117,118 (1978)), PSC-833 (cyclosporin D, 6-[(2S, 4R, 6E)-4-methyl-2-(methylamino)-3-oxo-6-octenoic acid]-(9CI)), [US Patent 5,525,590] [ACS 121584-18-7], Keller et al., Int J Cancer 50:593-597 (1992)), RU-486 (17\beta-hydroxy-11\beta-[4-dimethylaminophenyl]-17α prop-1-ynyl estra-4, 9-dien-3 one), RU-49953 (11β, 17α di[4-(dimethylamino)phenyl[estra-1,3,5(10)-triene-3, 17β -diol $\frac{17\beta}{\rho}$ -hydroxy- $\frac{11\beta}{\rho}$, $\frac{17\alpha-[4-1]}{\rho}$ dimethylaminophenyl] - 17α prop 1 ynyl estra 4, 9 dien 3 one), S9778 (6-{4-[2,2-di()ethylamino]-1-piperidinyl}-N,N', di-2-propenyl-1,3,5-triazine-2,4-diamine, sulfonate, [US patent 5,225,411; EP 466586] [ACS # 140945-01-3]; Dhainaut et al., "New triazine derivatives as potent modulators of multidrug resistance," J Medicinal Chemistry *35*:2481-2496 (1992)),MS-209 (5-[3-[4-(2,2-diphenylacetyl)piperazin-1-yl]-2hydroxypropoxylquinoline sesquifumarate, [US patent 5,405,843 (continuation of 5,112,817)], [ACS # 158681-49-3], Sato et al., "Reversal of multidrug resistance by a novel quinoline derivative, MS-209, Cancer Chemother Pharmacol 35:271-277 (1995)), MS-073 (Fukazawa et al., European Patent Application 0363212 (1989)), FK-506 (Tanaka et al., M. Physicochemical properties of FK-506, a novel immunosuppressant isolated from Streptomyces tsukubaensis" Transplantation Proceedings. 19(5 Suppl 6):11-6, (1987); Naito et al., "Reversal of multidrug resistance by an immunosuppressive agent FK-506," Cancer Chemother & Pharmacol. 29:195-200 (1992); Pourtier-Manzanedo et al., "FK-506 (fujimycin) reverses the multidrug resistance of tumor cells in vitro," Anti-Cancer Drugs 2:279-83 (1991); Epand & Epand, "The new potent immunosuppressant FK-506 reverses multidrug resistance in Chinese hamster ovary cells," Anti-Cancer Drug Design 6:189-93 (1991)), VX-710 (2-peperidinecarboxylic acid, 1-[oxo(3,4,5trimethoxyphenyl)acetyl]-3-(3-pyridinyl)-1-[3-(3-pyridinyl)propyl]butyl ester [ACS 159997-94-

1] [US patent number 5,620,971] Germann et al., "Chemosensitization and drug accumulation effects of VX-710, verapamil, cyclosporin A, MS-209 and GF120918 in multidrug resistanceassociated protein MRP" Anti-Cancer Drugs 8:41-155 (1997); Germann et al., "Cellular and biochemical characterization of VX-710 as a chemosensitizer: reversal of P-glycoproteinmediated multidrug resistance in vitro" Anti-Cancer Drugs 8:125-140 (1997)), VX-853 ([US patent number 5,543,423] [ACS # 190454-58-1), AHC-52 (methyl 2-(N-benzyl-Nmethylamino)ethyl-2, 6-dimethyl-4-(2-isopropylpyrazolo[1,5-a]pyridine-3-yl)-1,4dihyropyridine-3,5-dicarboxylate; [Japanese Patent 63-135381; European Patent 0270926] [ACS 119666-09-0] Shinoda et al., "In vivo circumvention of vincristine resistance in mice with P388 leukemia using a novel compound, AHC-52," Cancer Res 49:1722-6 (1989)), GF-120918 (9,10dihydro-5-methoxy-9-oxo-N-[4-[2-(1,2,3,4-tetrahydro-6,7-dimethoxyisoquinol-2-yl) ethyl]phenyl]-4 acridinecarboxamide,[US patent 5,604,237] [ACS # 143664-11-3] Hyafil et al., "In vitro and in vivo reversal of multidrug resistance by GF120918, an acridonecarboxamide derivative," Cancer Res 53:4595-4602 (1993)), and XR-9051 (3-[(3Z, 6Z)-6-Benzylidene-1methyl-2,5-dioxopiperazin-3-ylidenemethyl]-N-[4-[2-(6,7-dimethoxy-1,2,3,4tetrahydroisoquinolin-2-yl)ethyl]phenyl]benzamide hydrochloride, [ACS#57-22-7]).